SOP#: RPS-21 Establishing a Genomic Data Sharing Project and

Required Documents

Version #: 3.0 Next Review Date: 04/2024

Approved Date: 04/2022 Review Interval Period: Biennial

NCI Clinical Director Signature:

POLICY

The Genomic Data Sharing (GDS) Policy applies to all NIH intramural research that generates large-scale human or non-human genomic data as well as the use of these data for subsequent research. Large-scale data include genome-wide association studies (GWAS), single nucleotide polymorphisms (SNP) arrays, and genome sequence, transcriptomic, metagenomic, epigenomic, and gene expression data. See Appendix A for NCI Center for Cancer Research guidance on thresholds for required data sharing.

For studies that will generate large-scale genomic data, investigators are required to develop the following documents **PRIOR** to start the research:

Human Studies:

- 1) Institutional Certification Memo AND
- 2) Genomic Data Sharing Plan

Non-Human Studies:

1) Genomic Data Sharing Plan

An Institutional Certification Memo certifies that:

- data submission and sharing are consistent with the informed consent of the study participants
- consideration was given to risks to individual participants and their families associated with the shared data
- to the extent possible, consideration was given to risks, groups, or populations associated with the shared data
- the principal investigator's plans of de-identifying data sets are consistent with the GDS policy

More than one Institutional Certification Memo is required if the research began prior to August 31, 2015 (with or without consent) and continued past the policy implementation date when consent is mandatory. One memo is required for each instance:

- 1) prior to August 31, 2015, with consent
- 2) prior to August 31, 2015, without consent
- 3) after August 31, 2015, with consent

Effective November 1, 2018, the NIH Genomic Data Sharing Policy was expanded to enable **unrestricted access** to Genomic Summary Results (GSR). GSR are defined as any systematically computed statistics such as, but not limited to, genotype counts, and frequencies and allele counts and frequencies. The GSR are available through unrestricted access except when a population is determined to meet sensitive criteria. Sensitivity is defined as study populations from isolated geographic regions, or with rare or potentially stigmatizing traits which could result in increased privacy or confidentiality risks. The GSR committee is responsible for making the sensitivity determination with input from the investigators. If a sensitivity determination has not yet been made by the committee, contact the CCR GPA Kathleen Calzone.

Multi-Institutional studies in which NIH is the coordinating site and will be responsible for data sharing requires an Institutional Certification Memo from each site that is accruing samples and/or research participants unless NIH is the single IRB of record.

SOP#: RPS-21, v3.0

A **Genomic Data Sharing Plan** provides information on the proposed research that will generate large-scale human and non-human genomic data for which the GDS policy applies.

PURPOSE

The purpose of this standard operating procedure is to provide instructions for creating a GDS Project and corresponding Genomic Data Sharing Plan and, for human studies, the Institutional Certification Memo(s).

RESOURCES

- CCR Genomic Data Sharing Policy Website
- CCR Genomic Data Sharing submission portal
- NCI Genetic Data Sharing Policy <u>Website</u>
- NIH Genomic Data Sharing Policy
- NIH Genomic Data Sharing Website
- NIH Intramural Investigator GDS Responsibilities
- GDS Data Repositories and Trusted Partners

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PROCEDURES

Please have key protocol or study documents readily available to streamline the process.

Step 1: Create new Project, Enter all relevant Information

- All projects (human and non-human) that require compliance with the NIH GDS policy require basic project information.
 - Please refer to the CCR GDS website or contact the CCR GPA if you are uncertain whether your project/protocol falls within the purview of the GDS policy.
- Project Information is entered into the Genomic Data Sharing Portal
 - Sign into portal using your PIV card or NIH Username/password
 - o Institutional Certification Memo(s) and Genomic Data Sharing Plan are generated via the portal
- Required Project Information includes:
 - Project Title and Abbreviated Project title

- o Principal Investigator name
 - For clinical trials, Principal investigator is considered the protocol Principal Investigator or Lead Associate Investigator. For laboratory-based studies, Principal Investigator is considered the scientist leading the research.
- Email, phone number, title, and branch will be auto-populated from NIH Enterprise Directory (NED)
 - If there is an error, please contact your Administrative Officer to get NED content updated.
- Specify the organism type (human, non-human, or both)
- o Protocol ID
 - For clinical trials, please include the Protocol ID from iRIS. For laboratory-based studies please include a study identifier OR select N/A.
- Indicate the name/study number of the Parent Protocol/Project
 - Parent Protocols/Projects may be Tissue Procurement studies, OHRSP exemptions, or other umbrella projects in which multiple protocols/projects with different investigators are conducted under the umbrella protocol/project.
- Study Title and Abbreviated title
 - These may be different if your study is part of a larger project or the same as the Project title and can be copied and pasted. Please provide the same abbreviated Study title used in your protocol.
- If applicable: ZIA number, Journal mandate, Lead investigator, associated Study IDs

NOTE: Select Submit and please refresh the page before creating an IC memo or GDS Plan.

Step 2: Establishing Collaborator Access

You have the option of giving other NIH staff permission to access the Project and associated forms.

- Go to the User Access ribbon and click NEW
 - Enter the NIH User Name (which will link to full name and email)
 - Access type (Read Only or Read/Write)

Step 3: Select Forms to Complete

- Human Studies require a Institutional Certification Memo(s) AND Genomic Data Sharing Plan
- Non-Human Studies require a Genomic Data Sharing Plan only

Step 4: Institutional Certification Memo(s)

- An Institutional Certification Memo is established by clicking the Institutional Certification tab at the bottom of the Project page.
 - Note: more than one Institutional Certification Memo is required if the research began prior to August 31, 2015 (with or without consent) and continued past the policy implementation date when consent is mandatory. One memo is required for each instance:
 - 1) prior to August 31, 2015, with consent
 - 2) prior to August 31, 2015, without consent
 - 3) after August 31, 2015, with consent

- Only ONE Institutional Certification Memo can be generated at a time. If your project requires more than one memo, you will need to repeat the procedures described below for EACH instance requiring an Institutional Certification Memo.
- Click Create a NEW IC to start generating an Institutional Certification Memo.
- You must select at least one of the options below to generate the fields for a new IC memo.
 - Indicate if the samples have been collected before August 31, 2015 (Yes/No)
 - Indicate if the samples were collected with or without consent (Yes/No)
- Complete the starred information that is not already pre-populated from the Project Information
 NOTE: This portal may be used by other NIH Institutes so not all information provided is
 prepopulated for NCI.
 - Certification Date
 - Name of Institution NCI
 - Organization of GPA NCI
 - Organization of SD (Scientific Directors) NCI
 - o Original Study Name
 - Project Title for data to be submitted will be auto populated from the Project Information entered in Step 1.
 - Select data access, unrestricted or controlled access
 - Human studies are controlled access

If the release of Genomic Summary Results (GSR) was determined by the committee to be sensitive, check the controlled-access box (see figure below) AND provide an explanation in the box provided (see figure below)



NOTE: The GSR are available through unrestricted access **except** when a population is determined to meet sensitive criteria.

Add consent groups for each research site

SOP#: RPS-21, v3.0

NOTE: You must Select Submit prior to adding Consent Groups. This will establish your IC memo in the system and allow you to add Consent groups.

- Select New
 - Enter research site name (one site per row)
 - Select Data Use Limitations
 - Data Use Limitations are based on the terms of the informed consent of the study participants from whom the genomic data have been generated.
 - Click the ? to open definitions for each option.
 - Data Use Limitation Modifiers
 - These modifiers are also based on the consent.
 - Click the ? to open definitions for each option if applicable.
- Select Submit this will create the GDSP in the system you can then select one of the following:
 - Save enables you to return at a later time and edit content
 - Submit circulates the memo for review and approval/signature
 - Cancel all content entered will be withdrawn
- NOTE: You may have to complete more than one Institutional Certification Memo

Step 5: Genomic Data Sharing Plan

- A Genomic Data Sharing (GDS) Plan can be established by clicking the Genomic Data Sharing Plan tab at the bottom of the Project page.
- Click tab and select NEW to start generating a GDS Plan.
- Elements of a GDS Plan include the following:
 - Principal Investigator information, auto populated from the Project Information
 - Plan Submission Date
 - Tab 1: Project information
 - Tab 2: Data Source and type of samples (Check all that apply)
 - Tab 3: Type Select detailed information that applies to your GDS plan. Note: Some fields require # of samples once selected.
 - Tab 4: Submission Information these reflect any data use limitations specified in the consent.

Step 6: Review and Approval

- GPA review and approval
 - Submitted IC Memos and GDS Plans are first routed to the GPA for review.
 - Memos that require revision will be returned to the investigator with instructions from the GPA of the needed modifications.
 - Once the revisions have been made, resubmit the IC Memo or GDS Plan for review and approval by the GPA.
 - The submitter will be notified of GPA approval via an email generated from the submission system.

- Scientific Director review and approval
 - o GPA will submit the GPA approved IC memo or GDS Plan to the CCR Scientific Director for review and approval.
 - Approval by the Scientific Director completes the IC Memo or GDS Plan submission and approval process. The submitter will be notified, via email, of approval by Scientific Director.
 - For clinical studies, all completed GDS Plans will be available to the Protocol Support Office (PSO) staff. The PSO will upload a copy of completed Genomic Data Sharing Plan to the appropriate protocol specific regulatory file.
 - o For plans not approved by the Scientific Director:
 - GPA will work with the investigator to fulfill the requirements of the CCR policy.
 - Once a revised IC memo or GDS Plan is in place, the revised document will require resubmission for approvals.
- Once approved, the Institutional Certification memo and Genomic Data Sharing Plan may be viewed within your project. The Institutional Certification memo may be exported as needed.

SOP#: RPS-21, v3.0

Appendix A: NCI Guidance on Genomic Data Sharing

Examples of projects for which the NCI anticipates data sharing (*regardless of study design*) include, but are not limited to:

	# of Specimens	
	Human	Model Organisms,
	(including	Non-Human Cell
	human cell	Lines, Infectious
	lines)	Organisms
SNP array data from >500K single nucleotide polymorphisms	1,000	500
(SNPs) (e.g., GWAS data)		
DNA sequence data from < 100 genes or regions of interest	1,000	500
(e.g., targeted sequencing)		
DNA sequence data from ≥ 100 genes or regions of interest	100	50
(e.g., targeted sequencing, whole exome sequencing, whole		
genome sequencing)		
Genome-wide RNA sequencing (RNA-seq) data	100	50
(e.g., transcriptomic data)		
Genome-wide DNA methylation data	100	50
(e.g., bisulfite sequencing data)		
Genome-wide chromatin immunoprecipitation sequencing	100	50
(ChIP-seq) data		
(e.g. transcription factor ChIP-seq, histone modification ChIP-seq)		
Metagenome (or microbiome) sequencing data	100	50
(e.g., 16S rRNA sequencing, shotgun metagenomics, whole-		
genome microbial sequencing)		
Metatranscriptome sequencing data	100	50
(e.g., microbial/microbiome transcriptomics)		

<u>NOTE</u>: The number of samples includes distinct individuals, species, strains, samples, treatments, time points, and tissues. For example, data from 25 patients at four time points after treatment would reach a 100-sample threshold, as would data from 50 tumor-normal comparisons. Guidance on Genomic Data Sharing for rare diseases and rare cancers

Guidance issued from the Office of the Director, CCR mandates sharing of genomic data for projects examining rare diseases and rare cancers. There are **no minimum thresholds** to meet for such projects. The Trans-NCI Genomic Data Sharing Working Group has adopted the definition of rare disease – a disease that affects **less than 200,000 persons** in the United States, that has been <u>set forth</u> by the U.S. Food and Drug Administration (FDA).

Examples of smaller-scale projects that the NCI would likely mandate data sharing for include, but are not limited to:

- Projects examining rare cancers, rare-cancer-related outcomes, or rare cancer subtypes
- Projects focusing on under-studied populations

Examples of Research Outside the Scope of the GDS Policy:

Examples of NIH-funded research or research-related activities that are outside the Policy's scope include, but are not limited to, projects that do not meet the criteria in the above examples and involve:

- Instrument calibration exercises
- Statistical or technical methods development